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(54) Title: METHOD OF PREVENTING AGGREGATION AND COMPOSITIONS OBTAINED THEREI		VARIOUS SUBSTANCES UPON REHYDRATION OR THAWIN
(57) Abstract		
		variety of substances, (e.g. colloids, red blood cells, pharmaceutical whereby trehalose is added to a solution or suspension of the substance.
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METHOD OF PREVENTING AGGREGATION OF VARIOUS SUBSTANCES UPON REHYDRATION OR THAWING AND COMPOSITIONS OBTAINED THEREBY

5 Field of the Invention

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The present invention relates to methods of preventing the formation of aggregates of various substances upon dehydration and rehydration and upon freezing and thawing. Compositions obtained thereby are also encompassed by the 10 invention.

Background of the Invention

Storage and processing of a wide range of substances in a dehydrated or frozen form is necessary to retain 15 activity, prevent degradation products from forming and to facilitate handling and transport. Unfortunately, upon rehydration or thawing, many substances tend to aggregate, thereby decreasing their effective concentration and often rendering them useless or forming harmful byproducts.

Various methods have been tried to prevent or eliminate such aggregation. For instance, detergents and chaotropic agents are often used to prevent aggregation of proteins in solution. These agents are thought to prevent aggregation mediated by hydrophobic interactions and thus 25 are limited to prevention of aggregation due to this cause. See, e.g., Tanford and Reynolds (1976) Biochim. Biophys. Acta. 457:133; and Tanford, "The Hydrophobic Effect", 2nd · Ed., Wiley, N.Y. (1980). Such agents may also not be suitable for use where the substances are to be formulated 30 into therapeutic compositions as they may cause adverse reactions.

Aluminium salts in solution are in the form of a highly hydrated colloidal gel and carry a surface charge at any pH 35 outside their isoelectric point. Since each colloidal particle carries the same charge, they mutually repel each other and thus naturally form a stable colloidal gel. When

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the hydration shell is removed (e.g., by freezing or drying) the particles can contact each other and the surface energy causes aggregation.

Trehalose, α -D-glucopyranosyl- α -D-glucopyranoside, is a naturally occurring, non-reducing disaccharide which was initially found to be responsible for protection of intact plant cells from desiccation. Trehalose has been shown to be useful in preventing denaturation of proteins viruses and foodstuffs during desiccation. U.S. Patent

Nos. 4,891,319; 5,149,653; 5,026,566; Blakeley et al. (1990) Lancet 336:854-855; Roser (1991) Trends in Food Sci. and Tech. pp.166-169; Colaco et al. (1992) Biotechnol. Internat., pp. 345-350; Roser (1991) BioPharm. 4:47-53; and Colaco et al. (1992) Bio/Tech. 10:1007-1011.

In the field of protein purification it would be particularly useful to eliminate or prevent the tendency of, eg. proteins, to aggregate upon rehydration and thawing. This is especially important in the area of biopharmaceuticals where the proteins are often used as an ongoing basis of treatment. In the case where protein aggregates form and are injected into a patient, antibodies may form to the protein which diminish the effectiveness of the treatment. Thus, it would be useful to prevent aggregation of a wide variety of substances particularly those useful in medicine.

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Summary of the Invention

The invention encompasses a method of reducing aggregation during dehydration and rehydration of substances comprising the steps of adding to a solution or suspension of the substances an amount of trehalose sufficient to prevent aggregation upon rehydration; and dehydrating the solution or suspension. The invention also encompasses the compositions obtained thereby.

The invention further encompasses rehydrating the
solution or suspension to obtain a composition
substantially lacking aggregates of the substance. The
compositions obtained thereby are also encompassed by the
invention.

The invention further encompasses a method of reducing
aggregation of substances in solution or suspension during
freezing (and optionally thawing) comprising the steps of
adding to the solution or suspension of the substance an
amount of trehalose sufficient to prevent aggregation
during freezing; and freezing the solution or suspension.
The invention also comprises the compositions obtained
thereby.

The invention further comprises the step of thawing the frozen solution or suspension to obtain a composition substantially lacking aggregates of the substance. The compositions obtained thereby are also encompassed by the invention.

A wide variety of substances are suitable for use in the invention including, but not limited to, therapeutic, prophylactic and diagnostic.

When the substance is red blood cells, the method may further comprise the step of fixing the red blood cells prior to adding trehalose. Fixing of red blood cells can be done by any known method including, but not limited to, formaldehyde and glutaraldehyde.

Brief Description of the Drawings

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Figure 1 is a bar graph depicting the percent height of sedimentation of aluminium phosphate per column after 24 hours. Columns labeled with + and - symbols were dried in the presence and absence of trehalose respectively. 5 stands for vacuum drying, Tfd stands for freeze drying, Tfz strands for freezing, T4fz stands for freeze thawing four times and Tw stands for aqueous samples.

Figure 2 is a bar graph depicting the percent height of sedimentation of aluminium phosphate per column after 10 5.5 hours. Prior to testing, samples were stored for one week at 45°C. The abbreviations are the same as those in Figure 1.

Figure 3 is a bar graph depicting the percent height of sedimentation of aluminium hydroxide after 24 hours. 15 The numbers refer to the series as described in Example 3, d stands for vacuum drying, w stands for aqueous control, and f stands for freezing.

Detailed Description of the Invention

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The present invention encompasses a method of reducing aggregation during dehydration and rehydration of substances by adding to an eg. aqueous solution or suspension of the substances an amount of trehalose sufficient to prevent aggregation upon rehydration; and 25 dehydrating the solution or suspension.

The invention further encompasses a method of reducing aggregation of substances in an eg. aqueous solution or suspension during freezing and thawing comprising the steps of adding to the solution or suspension of the substance an 30 amount of trehalose sufficient to prevent aggregation during freezing and thawing; and freezing the solution or suspension.

The invention also relates to eg. aqueous compositions containing trehalose in order to prevent or reduce 35 aggreagtion, including frozen and dehydrated compositions.

The term "aggregation" as used herein refers to the interaction of two or more molecules of a substance such

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that they no longer behave as monomers but as dimers, trimers or other multimeric forms. Reducing aggregation decreases the concentration of multimeric forms compared to substances dehydrated and rehydrated or frozen and thawed in the absence of trehalose. A substance substantially free of aggregates or substantially nonaggregated is one which, upon rehydration or thawing, contains a decreased amount of multimeric forms of the substance compared to a control lacking trehalose. Typically, trehalose prevents the formation of all multimeric forms of the substance. In the case of growth hormone, for instance, the addition of trehalose prior to dehydrating or freezing can result in the elimination of all multimeric forms with the exception of dimers. The dimers are, however, reduced in comparison to a control.

In a preferred embodiment, the substances suitable for use in the invention have medical utility. Such substances include, but are not limited to, therapeutic substances, prophylactic substances and diagnostic substances. The substances can be those which form multimers upon dehydration/rehydration and/or freezing/thawing. The method of formation of multimers or aggregates is not critical to the invention.

Suitable therapeutic substances include, but are not
limited to, any therapeutically effective biological
modifier. Such modifiers include, but are not limited to,
proteins and peptides, steroid hormones, oligosaccharides,
nucleic acids, polynucleotides and a variety of small
molecules. Further, the modifiers may be derived from
natural sources made by recombinant or synthetic means and
include analogues, agonists and homologs. As used herein
"protein" refers also to peptides and polypeptides. Such
proteins include, but are not limited to, growth hormones,
growth factors, insulin, monoclonal antibodies, interferons
and interleukins. Preferably, the growth hormone is human
growth hormone. Suitable steroid hormones include, but are
not limited to, estrogen, progesterone and testosterone.

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Therapeutic substances prepared by the methods described herein are also encompassed by the invention.

Suitable prophylactic substances include, but are not limited to, aluminium based adjuvants such as aluminium 5 hydroxide and aluminium phosphate which are used in preparation of vaccines. Compositions containing the, eg. prophylactic, substances are further encompassed by the invention. Preferred compositions include vaccines containing the aluminium hydroxide or aluminium phosphate 10 prepared by the method described herein. Suitable vaccines include, but are not limited to, combination vaccines, such as diphtheria, tetanus, pertussis (DTP) or DTP/inactivated poliovaccine (IPV). Suitable diagnostic substances include, but are not limited to, colloidal gold, polystyrene latex, fixed erythrocytes and monoclonal antibodies. Diagnostic substances and compositions prepared by the method described herein are also encompassed by the invention.

The dehydration step can be performed by any method
known in the art including, but not limited to,
lyophilization, drying at ambient conditions or drying
under reduced vapor pressure. When drying at reduced vapor
pressure, the temperature at which the drying occurs is
preferably below the temperature at which degradation of
the substance occurs.

The freezing step can be performed by any method known in the art including, but not limited to immersing in liquid nitrogen, placing in a freezer which may be at -4°C to -80°C, dry ice and alcohol freezing bath. The samples should be maintained at a temperature suitable to maintain the frozen state. Thawing the frozen sample can be by any means known in the art, for instance at room temperature or at an elevated temperature. If thawing occurs at an elevated temperature, the temperature should be below that which causes denaturation or other chemical changes in the substance. Optimal freezing and thawing temperatures can

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be determined empirically. Such a determination is within the skill of one in the art.

Once the substances have been dehydrated or frozen, they can be stored indefinitely. The dehydrated substances can store well at ambient temperatures, although they may be stored at any temperature below that which causes denaturation or other chemical changes. The invention further includes rehydration of the dehydrated samples to obtain solutions and suspensions substantially free of aggregates of the substance. Rehydration may add at least an amount of water sufficient to restore the buffer composition of the original solution or suspension but may add any amount of water or buffer.

When the substance comprises red blood cells, the

method may further comprise fixing the red blood cells

prior to adding trehalose. The fixing step may be done by

any method known in the art including, but not limited to,

glutaraldehyde. In the preferred embodiment, the cells are

fixed.

The methods of the present invention prefer that the trehalose be present in an amount sufficient to prevent or reduce aggregation of the substance upon rehydration or thawing. Such a determination will be made empirically and is well within the skill of one in the art. Preferably,

25 trehalose is added in an amount to obtain a final concentration of from about 1% to 50% (w/v). More preferably, trehalose is added in an amount to obtain a final concentration of from about 5% to 25% (w/v).

Trehalose is available from a variety of suppliers.

30 Preferably the grade of trehalose used is ANALAR reagent, molecular biology or pharmaceutical grade. In the case of medicinal compositions the trehalose preferably meets the good manufacturing practice (GMP) standards set by the Food and Drug Administration (FDA).

35 The invention also encompasses the products obtained by the method both before and after rehydration or thawing. In one embodiment, the invention includes the frozen

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compositions containing a substance and an amount of trehalose sufficient to prevent aggregation of the substance upon thawing. In another embodiment, the invention includes a dehydrated composition comprising a substance and an amount of trehalose sufficient to prevent aggregation of the substance upon rehydration. The invention further includes the compositions after being thawed or rehydrated respectively.

The invention additionally encompasses the use of trehalose as an aggregation reducing or preventing agent, particularly for substances present in an (eg. aqueous) composition such as a solution or suspension of that substance.

Interestingly, the amount of trehalose found to be
effective at preventing aggregation cannot be directly
extrapolated from the amount of trehalose effective in
preventing desiccation damage. For instance, work
presented in United States Patent No. 4,891,319 showed that
amounts of trehalose as low as 1% w/v in a protein solution
could prevent desiccation damage to proteins such as Factor
VIII. The Examples presented herein show that more than
30% w/v trehalose is necessary to completely prevent
aggregation of aluminium hydroxide and 15% w/v is necessary
to prevent aggregation of a protein.

The following examples are meant to illustrate, but not limit, the invention.

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EXAMPLES Example 1

<u>Prevention of Aggregation of Particulate</u> <u>Suspensions by Trehalose</u>

In order to determine whether trehalose prevented aggregation of particulate suspensions, two examples, colloidal gold and polystyrene latex, were studied. Colloidal gold was obtained from the Babraham Laboratories and polystyrene latex was a suspension of particles of polystyrene which had been purchased from Sigma Chemical Company.

The colloidal gold was made according to the method described by Frens (1993) Nature 241:20. It was dried from a concentrated suspension of 0.2% Au in a volume of 50 µl per well in a 96 well microtiter plate either with added 10% w/v trehalose or without trehalose and subsequently rehydrated after storage for one week at 37°C in a dry oven. On rehydration, the material that had been dried in the presence of trehalose gave a smooth suspension of colloidal gold as determined by microscopic examination. The material that had been dried without trehalose showed microscopic aggregates which could not be broken up into a smooth suspension.

With the polystyrene latex, similar experiments were conducted. The latex was obtained from Sigma Chemical Company catalogue number LB-8, average diameter 0.8 micron polystyrene. It was used at the concentration obtained from the supplier and again was dried without any addition of, and also with the addition of, 10% w/v trehalose which was dissolved in the solution before drying. Both samples were rehydrated about a week after drying and were stored at 37°C in a dry oven in the interim. The material dried without trehalose was badly aggregated into very large clumps. The material dried in the presence of trehalose resuspended into a very smooth, single particulate suspension.

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Thus, the addition of trehalose prior to drying the particulate suspensions substantially reduced the amount of aggregation upon rehydration compared to a control lacking trehalose.

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Example 2 Effect of Trehalose-Drying on Aggregation of Red Blood Cells

10 Experiment

Rat RBCs were washed three times in an anti-coagulant CPD (102 mM trisodium citrate, 1.08 mM sodium phosphate and 11 mM dextrose), filtered through cotton wool and fixed in either 1% formaldehyde or 0.5% glutaraldehyde. Fixing was at room temperature for one hour. The fixed cells were washed three times in CPD and resuspended in either 10% trehalose and 0.12 mM sodium azide (NaN₃) or CPD. The final cell concentration was 25% w/v.

Cells fixed in formaldehyde lysed on washing and were 20 not processed further.

Unfixed cells agglutinated in trehalose and needed the addition of 1/5th volume of phosphate-buffered saline before being processed further.

Then 100 μ l of cells in either 10% trehalose 0.12 mM NaN₃ or CPD were dried either in Nunc plates or on slides and examined microscopically for aggregates.

Results

The unfixed cells dried without trehalose lysed

30 completely and with those dried with trehalose also showed

95-99% lysis though the ghosts showed discoid morphology.

The fixed cells dried without trehalose showed gross macroscopic aggregation of the cells. The fixed cells dried with trehalose resuspended as a smooth single cell suspension with only a few microaggregates. These microaggregates appear to form at higher concentrations of

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trehalose and thus do not appear to be concentration dependent.

Example 3

Aggregation of Aluminium Hydroxide and Phosphate

Sedimentation Assay

The following method was followed to determine whether trehalose was successful in preventing aggregation of prophylactic adjuvants.

Aluminium phosphate and aluminium hydroxide were diluted 5-fold to a final concentration of 0.6% w/v and allowed to sediment in 1 ml glass pipettes. The height of the sediment column was measured at various time intervals up to 24 hours. Note that the % height of sediment column should not be < 30% when a steady state has been reached (about 5 hours.)

The samples were dried under vacuum, frozen at $-20\,^{\circ}\text{C}$ and thawed at room temperature.

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Results

<u>Pilot 1</u>. Aluminium phosphate

Different forms of drying and storage were compared in the presence or absence of 15% trehalose. These were vacuum drying (Tv), freeze drying (Tfd), freezing (Tfz) and freeze thawing for four cycles (T4fz). Wet controls (Tw) which were stored at 4°C were also run.

Then 200 μ l sample per glass vial were dried and sedimentation assays carried out at day 0 and after 1 week storage at 45°C. The results obtained are shown in Figures 1 and 2.

Pilot 2. The aggregation of aluminium hydroxide and haemaccel (degraded gelatin) was measured with a titration of trehalose with the concentrations shown in Table 1. The samples contained 1.5% aluminium hydroxide and 2% haemaccel. Only vacuum drying (d) and freezing (f) were compared. Wet controls (w) contained trehalose and

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haemaccel but were not dried or frozen. Each series contained (d), (f) and (w) samples. The concentrations used are shown in Table 1 and the results obtained are shown in Figure 3.

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		<u>Table 1</u>	
	<u>Series</u>	Final % trehalose	<pre>% haemaccel</pre>
	1	7.5	_
	2	15	-
10	3	30	<u>.</u>
	4	15	2
	5	<u>-</u>	_

Conclusions

- a) 15% trehalose can prevent freezing induced aggregation in aluminium phosphate and aluminium hydroxide
 - b) 7.5% trehalose is not sufficient for preventing aggregation during the drying process.
- 20 c) No additional effect of Haemaccel at 2% was observed.

Aluminium hydroxide, dried in the absence of trehalose and rehydrated was found to be aggregated into large clumps which sedimented rapidly and quickly to yield a very small gel column. Trehalose in concentrations above 15% prevented this aggregation so that the rehydrated material formed a gel column of a height similar to the fresh, undehydrated material. This sedimentation pattern illustrates that the hydrated, nonaggregated molecules have a large hydration shell volume and are separated from one another causing them to sediment slowly.

Example 4

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Effect of Trehalose on

Aggregation of Biological Molecules

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Protein formulations may undergo modification by a number of mechanisms including deamidation, oxidation and aggregation, the principle causes of human growth hormone (hGH) degradation. Deamidation and oxidation are considered collectively as chemical degradation. To date there is little evidence of any effect of these chemical degradation products on biopotency. Pearlman and Bewly (1993) In: Wang and Pearlman eds. Stability and Characterization of Protein and Peptide Drugs, pp. 1-58, Plenum Press, New York.

Aggregation is the principle problem affecting hGH and other protein formulations used as biopharmaceuticals and may reduce biopotency. Soluble or insoluble aggregates can form as a result of both covalent and non-covalent interactions. A variety of stresses such as heating, freezing or agitation may induce aggregation. Whilst a visible insoluble aggregate may render a parenteral product unuseable, the major problem is the induction of an unwelcome immune response in the subject (Pearlman and Bewley, 1993). This is particularly detrimental where the protein formulations such as hGH are administered parenterally and in multiple doses.

The following experiment was performed to determine whether or not trehalose affected the aggregation of proteins. Samples of hGH (5 mg) were dried from 200 µl containing 15% trehalose, 5 mM Na₂HPO₄ - 2H₂O adjusted to pH 7.4 with H₃PO₄ (formulation A). Two control samples were prepared: 5 mg hGH dried from 200 µl sodium phosphate buffer pH 7.4 (formulation B); and 5 mg hGH dried from 200 µl sodium phosphate buffer pH 7.4, 5 mg glycine, 25 mg mannitol (formulation C). These formulations were dried for 20 hours in a vacuum drier at a pressure of 30 millitorr and a shelf temperature of 40°C. They were subsequently sealed under vacuum in standard pharmaceutical serum vials with rubber closures and a crimped aluminium seal.

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Following storage at 40°C in a dry incubator, samples were rehydrated with deionised water and analysed by reverse phase and size exclusion high performance liquid chromatography to determine chemical degradation and aggregation respectively according to the method described by Pikal et al. (1991) Pharm. Res. 8:427-436. These results are presented in Table 2.

Formulation A was subsequently re-analysed and compared with a conventionally freeze-dried essentially as described in Pikal et al. (1991) equivalent formulation (formulation D). These results are presented in Table 3.

Results

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An accelerated aging protocol of four weeks at 40°C

15 was utilized to assess stability and aggregation. The
formulation containing trehalose performs very well under
these conditions. No chemical degradation was observed and
the limited aggregation detected was restricted to dimer
formulation (Table 2, lines 1-4). The absence of high

20 molecular weight aggregates is significant.

Two hGH controls were formulated, one without a stabilizing excipient (B) and one containing glycine and mannitol that was similar to commercial formulations (C) (Table 2, lines 5-6). These formulations suffered from considerable chemical degradation and aggregate formation, both dimer and higher molecular weight. The values for the glycine mannitol formulation were comparable with results from a previous study in which a similar formulation was freeze-dried (Table 3, line 7, Pikal, et al. 1991). When the stability of formulation A was compared with that of a freeze-dried equivalent (formulation D), no difference in terms of 40°C stability was observed (Table 3, lines 1-6). In Tables 2 and 3 chemical degradation is measured by the area under the curve represented by the deamidated protein.

Thus the hGH formulations containing trehalose, either dried at 40°C or freeze-dried, have been shown to be considerable improvements on previous formulations.

Table 2
Summary of hGH Stabilization and Aggregation Data

(Part 1)

- 15 -

5	Line	Formulation	Treatment	% Chemical Degradation	% Aggregation Dimer	% Aggregation High Mol. Weight
	· 1	. A	pre-dry	3.1	0.4	0.003
	2	A	post-dry	3.3	0.6	0.06
10	3	A	2wk.,40°C	3.5	0.9	0.02
	4	А	4wk.,40°C	3.4	1.1	0.002
	5	В	4wk.,40°C	11.1	6.9	2.1
	6	С	4wk.,40°C	8.2	2.2	0.8

Table 3

Summary of hGH Stabilization and Aggregation Data

(Part 2)

20	Line	Formulation	Treatment	% Chemical Degradation	% Aggregation
	1	A	initial	4.15	0.66
	2	A	2wk.,40°C	4.16	0.92
	3	A	4wk.,40°C	4.25	1.04
25	4	D	initial	4.05	0.71
	5	D	2wk.,40°C	4.09	0.86
	6	D	4wk.,40°C	4.17	0.92
	7	E	4wk.,40°C	8.2	3.0

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Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity and understanding it will be apparent to those skilled in the art that certain changes and modifications may be practiced. Therefore, the description and examples should not be construed as limiting on the invention, which is delineated by the appended claims.

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CLAIMS

 A method of reducing or preventing aggregation during dehydration and rehydration of a substance, the
 method comprising:

adding to a solution or suspension of the substance an amount of trehalose sufficient to prevent or reduce aggregation upon rehydration; and

dehydrating the solution or suspension.

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2. A method according to claim 1 further comprising rehydrating the substance to obtain a solution or suspension of the substance in a substantially nonaggregated form.

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3. A method of reducing or preventing aggregation of a substance in solution or suspension during freezing, the method comprising:

adding to the solution or suspension of the substance 20 an amount of trehalose sufficient to reduce or prevent aggregation during freezing; and

freezing the solution or suspension.

- 4. A method according to claim 3 further comprising thawing the solution or suspension to obtain a solution or suspension of the substance in a substantially nonaggregated form.
- 5. A method according to any preceding claim wherein the substance is a therapeutic, prophylactic or diagnostic substance.
- 6. A method according to claim 5 wherein the substance is a therapeutic substance and is a biological response modifier.

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7. A method according to claim 6 wherein the biological modifier is a protein, peptide, steroid hormone, oligosaccharide, nucleic acid, polynucleotide or a small molecule.

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- 8. A method according to claim 7 wherein the protein is a growth hormone, growth factor, insulin, monoclonal antibody, interleukin or interferon.
- 9. A method according to claim 8 wherein the substance is human growth hormone.
 - 10. A method according to claim 7 wherein the steroid hormone is estrogen, progesterone or testosterone.

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- 11. A method according to claim 5 wherein the substance is a prophylactic substance and is an aluminium based adjuvant.
- 12. A method according to claim 11 further comprising incorporating the adjuvant into a vaccine.
- 13. A method according to claim 12 wherein the
 vaccine is a diphtheria/tetanus/pertussis (DTP) or
 25 diphtheria/tetanus/pertussis/inactivated poliovaccine
 (DTP/IPV).
 - 14. The method according to claim 13 wherein the vaccine is DTP or IPV.

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15. A method according to any preceding claim wherein the substance is a diagnostic substance and is colloidal gold, a polystyrene latex, fixed erythrocytes or a monoclonal antibody.

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- 16. A method according to any preceding claim wherein the trehalose is added in an amount to obtain a final concentration of from 1% to 50% (w/v).
- 5 17. A method according to any preceding claim wherein the trehalose is added in an amount to obtain a final concentration of from 5% to 25% (w/v).
- 18. A method according to any of claims 15 to 17

 10 wherein the substance is red blood cells and the method further comprises fixing the red blood cells prior to adding trehalose.
- 19. A method according to claim 18 wherein the fixing is by glutaraldehyde.
 - 20. A method according to any of claims 1, 2 or 5 to 19 wherein dehydration occurs by lyophilization, drying at ambient conditions or drying under reduced vapor pressure.

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21. A method according to claim 20 wherein the drying at reduced vapor pressure occurs at room temperature or at a temperature elevated above room temperature but below a temperature at which degradation or chemical change of the substance occurs.

- 22. A composition obtained by a method according to any preceding claim.
- 23. An aqueous composition comprising a substance and an amount of trehalose sufficient to reduce or prevent substantial aggregation of the substance upon freezing and thawing or dehydrating and rehydrating.
- 35 24. A frozen composition comprising a substance and an amount of trehalose sufficient to reduce or prevent substantial aggregation of the substance upon thawing.

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- 25. A dehydrated composition comprising a substance and an amount of trehalose sufficient to reduce or prevent aggregation of the substance upon rehydration.
- 5 26. A composition according to any of claims 23 to 25 which is a therapeutic, proplylactic or diagnostic composition.
- 27. Use of trehalose to reduce or prevent aggregation 10 of a substance upon freezing and thawing or dehydrating and rehydrating in a solution or suspension of that substance.

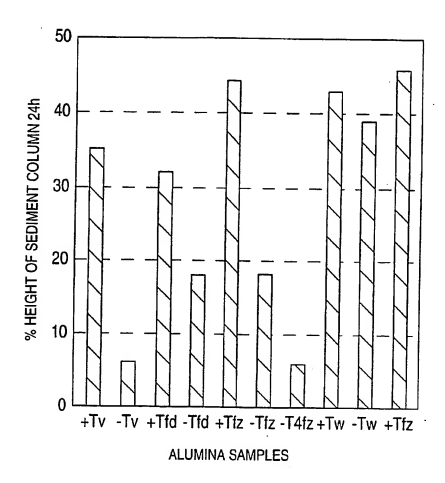


FIG. 1

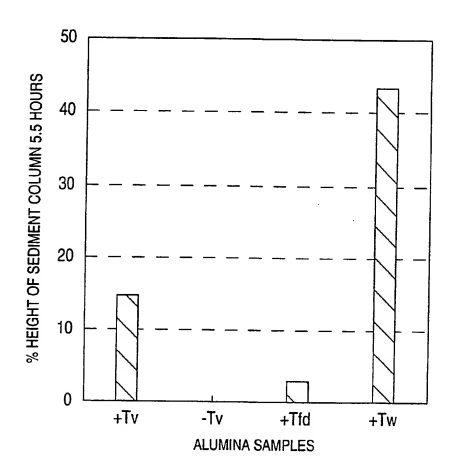


FIG. 2

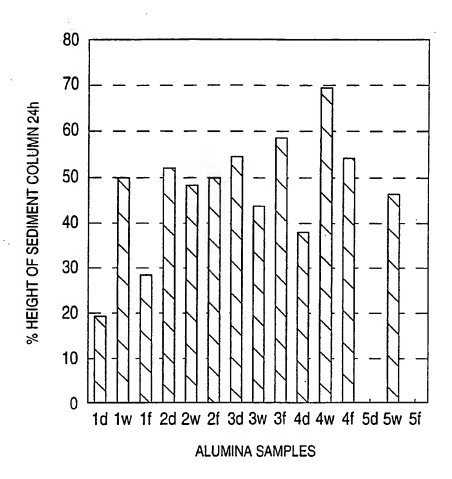


FIG. 3

INTERNATIONAL SEARCH REPORT

Internation Application No PCT/GB 95/01277

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 A61K47/26 C07K1/ C07K1/00 C12N9/96 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 A61K CO7K C12N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. X WO,A,90 04329 (COULTER CORP) 3 May 1990 1-27 see page 4, line 1 - line 29 see page 7, line 20 - line 30; claims 12-16 X WO,A,90 13637 (HYGEIA SCIENCES LTD) 15 1-27 November 1990 see page 16, line 20 - page 17, line 9 US,A,5 026 566 (ROSER BRUCE J) 25 June 1-27 cited in the application X see the whole document 22-26 Y US,A,4 891 319 (ROSER BRUCE J) 2 January 1-27 1990 cited in the application X see the whole document 22-26 -/--X Further documents are listed in the continuation of box C. Patent family members are listed in annex. * Special categories of cited documents: later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the document defining the general state of the art which is not considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) 'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-ments, such combination being obvious to a person skilled in the art. 'O' document referring to an oral disclosure, use, exhibition or document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 2 2, 11, 95 19 October 1995 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Riswijk Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+ 31-70) 340-3016 Foerster, W

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